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Paediatric Dentistry

Low level light therapy in the management of paediatric oral and oropharyngeal mucositis

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Abstract (100 words)

Oral and oropharyngeal mucositis is a common, debilitating condition experienced by patients undergoing oncology treatment. There are many different management strategies, with low level light therapy (LLLT) an emerging field.

Ongoing research on the topic of LLLT for mucositis has resulted in LLLT being included in national clinical guidelines. The number of centres currently using LLLT for paediatric patients with oropharyngeal mucositis is growing, with Glasgow having successfully used this treatment method for a number of years.

Across medicine and dentistry, LLLT is coming to the fore and is a treatment modality we should all be aware of.

CPD/Clinical Relevance

New technologies and treatment modalities are areas that practitioners should stay up to date with. Low level light therapy continues to be a growing research field.

Objective: The reader should understand the significance of oral and oropharyngeal mucositis for the paediatric oncology patient and the relevance of low level light therapy towards its management.

Oral and Oropharyngeal mucositis

Oral and oropharyngeal mucositis occurs in 52-80% of children undergoing treatment for cancer.¹ Mucositis is a debilitating condition which can be so severe it can delay or stop oncology treatment. The mucositis disease process is initiated by chemotherapy or radiotherapy at 0-2 days, followed by cell damage at 2-10 days. Frank ulceration and mucosal damage occurs at 10-15 days and finally healing after 2-3 weeks. The clinical presentation is a combination of erythema and ulceration; and as healing begins, mucosal sloughing affects the intra-oral tissues and oesophageal tract. Maintaining adequate oral hygiene is difficult due to oral discomfort which may further exacerbate the condition and increase a patient's vulnerability to developing infection. The consequences for paediatric patients are significant; eating, drinking, speaking and engaging with those around them can be significantly restricted which presents concerns regarding recovery and physical and mental wellbeing. Psychological distress has been described by parents of children suffering from mucositis at a time when stress and anxiety of the parent and child is already amplified by many other factors.²



Figure 1

Mucositis can be graded using the ‘World Health Organisation (WHO) Grading,’ which comprises gradings zero to four ³, as can be seen in the table below. Both the clinical presentation and functional outcomes form the basis for this grading system:

WHO Mucositis Grading

WHO Grading	Description
0	Healthy mucosa
1	Pain with no ulceration
2	Ulcers and erythema but the patient is still able to swallow a solid diet
3	The patient cannot swallow a solid diet;
4	Mucositis is so severe that alimentation is not possible

Management of mucositis

General management of mucositis often involves a mixture of preventive and symptomatic control measures such as; oral hygiene instruction, soft diet, cryotherapy, protective gels, analgesics ranging from paracetamol and non-steroidal analgesia to narcotic analgesia, topical anaesthetics, benzydamine hydrochloride sprays or mouthwashes and chlorhexidine and calcium phosphate mouthrinses. Many of these, for example mouthwashes, require a certain degree of cooperation and maturity which not all paediatric patients will have. Mucositis is such a common and significant problem in oncology that many different guidelines have been developed to assist practitioners. Guidelines published in 2012 by the The Royal College of Surgeons of England and The

British Society for Disability and Oral Health ⁴ recommend benzydamine hydrochloride mouthwash, cryotherapy, intravenous keratinocyte growth factor-1 (palifermin), anaesthetic mouthwash and analgesia as options to consider in the management of mucositis. More recent guidelines published in 2014 by the Multinational Association of Supportive Care in Cancer/International Society of Oral Oncology (MASCC/ISOO) similarly advises most of the above mentioned interventions along with other weaker evidence for options such as doxepin mouthwash and zinc supplements.⁵ A guideline by Sung et al in 2015 included a weak recommendation for cryotherapy to prevent mucositis.⁶ Both MASCC/ISOO ⁵ and Sung et al ⁶ also recommend LLLT as an option for managing mucositis. It is anticipated with new evidence emerging that all future guideline revisions will also soon advocate Low Level Light Therapy.

Low Level Light therapy

Low level light therapy (LLLT) is emerging as an effective treatment modality in both medicine and dentistry for a range of conditions. LLLT involves the use of a low power laser emitting less than 500mW on the red or near infra-red spectrum with light wavelengths of 600nm-1000nm. On applying the light to areas of the body, cells are stimulated thereby reducing inflammation, increasing cell metabolism and inducing endorphins. A Dental Update article by Milward et al. in 2014 ⁷ describes in further detail the mechanism of action including wound repair, analgesia and anti-inflammatory effects. The combination of cellular effects has led to this therapy becoming a widely studied topic and gathering increasing attention from the medical and dental professions. Despite the growing body of evidence, a recent systematic review of by Hadis et al ⁸ found inadequacies in the reporting of LLLT studies, with incomplete information available on wavelength, light source type, power, pulse frequency, beam area, irradiance, exposure time, radiant energy and fluence

As mentioned, low level light therapy can be considered for a number of different conditions. Research in medical patients has resulted in a Cochrane review on the use of LLLT in rheumatoid arthritis, providing evidence that it reduces pain and stiffness.⁹ Systematic reviews on its use in treating trigeminal neuralgia¹⁰ and recurrent aphthous ulceration¹¹ are also available in the literature. Ayyildiz et al discusses LLLT for use on patients suffering from Temporomandibular Joint Dysfunction with limited mouth opening.¹² There has also been research involving orthodontic patients concerning the potential for LLLT to accelerate tooth movement and also reduce orthodontic related pain^{13,14}.

Low Level Light therapy and mucositis

In the literature there have been many studies involving LLLT as a treatment modality for oral and oro-pharyngeal mucositis. In a systematic review with meta-analysis by Bjordal et al,¹⁵ Low Level Light Therapy was shown to reduce pain, severity and duration of mucositis. This paper included 11 randomised placebo controlled trials with a total of 415 patients. The relative risk of developing mucositis was found to be significantly reduced after LLLT when compared to placebo (RR = 2.03 p value = 0.02). In patients with mucositis, the number of days patients had mucositis was found to be reduced for severity gradings 2 or more from 4.38 days to 1.33 days compared with placebo.

A randomised controlled study by Khun et al¹⁶ in 2009, recommended LLLT as a first-line option in paediatric patients with chemotherapy-induced oropharyngeal mucositis. The trial compared treatment for mucositis with placebo for 5 days compared to LLLT for the same duration of time. Twenty-one patients were included in the study with the number of days

patients had mucositis decreasing from a mean duration of 8.9 days with placebo to a mean duration of 5.8 days with LLLT.

The benefits that LLLT can offer to the paediatric patient appear to be significant. As this is often a patient group where cooperation may be limited, quick, efficient and effective treatment is necessary. In addition paediatric oncology patients in particular are a group where the prevention of infection and pain are of paramount importance.

Low Level Light therapy use for paediatric patient in Glasgow

Currently in the Royal Hospital for Children (previously The Royal Hospital for Sick Children) in Glasgow, LLLT is used daily in the management of oro-pharyngeal mucositis by the Paediatric Dentistry department. It is used often in conjunction with previously described measures such as mouth rinses, analgesia and protective gels. The paediatric dental team makes regular visits to the oncology ward and has a good relationship with the ward staff allowing a team approach to the provision of LLLT. LLLT is delivered using a Diobeam 830 laser (830nm) with an output of 150mW (see Figure 2). Bensadoun et al 2015¹⁷ recommended that wavelengths of 633-685nm or 780-830 nm and power output between 10 and 150 mW are used to treat mucositis.

Figure 2



Initially an extra- and intra-oral examination by a qualified dentist determines the areas affected by mucositis and a WHO mucositis grading is assigned. An administered dose (J/cm²) is then delivered according to the severity of mucositis.³ The paperwork completed for each treatment episode used can be seen below in Figure 3. Although in Glasgow the 'laser' is primarily used for symptomatic mucositis, one joule setting per centimetre squared is available for use on healthy mucosa, which would allow for the laser to be given as prophylactic treatment. The Diobeam laser is classed as a 3B laser which means it carries an ocular hazard. Local laser safety precautions are employed, such as using safety glasses to protect the patient, parent's and user's eyes. The laser is used in an approved room which has blinds on the windows, a locked door and safety signs displayed outside. LLLT is not visible but has a red guide light to help the user to visualise the target area and an auditory guide for when the laser is active. Treatment is painless. Similar to radiographic regulations, there are also local laser rules which have to be followed and training must be completed prior to any clinician delivering treatment.

Figure 3

Oral Mucositis – treatment proforma

Diagnosis

Name CHI..... Date

Neutrophils

Date of first presentation of mucositis

Duration of mucositis

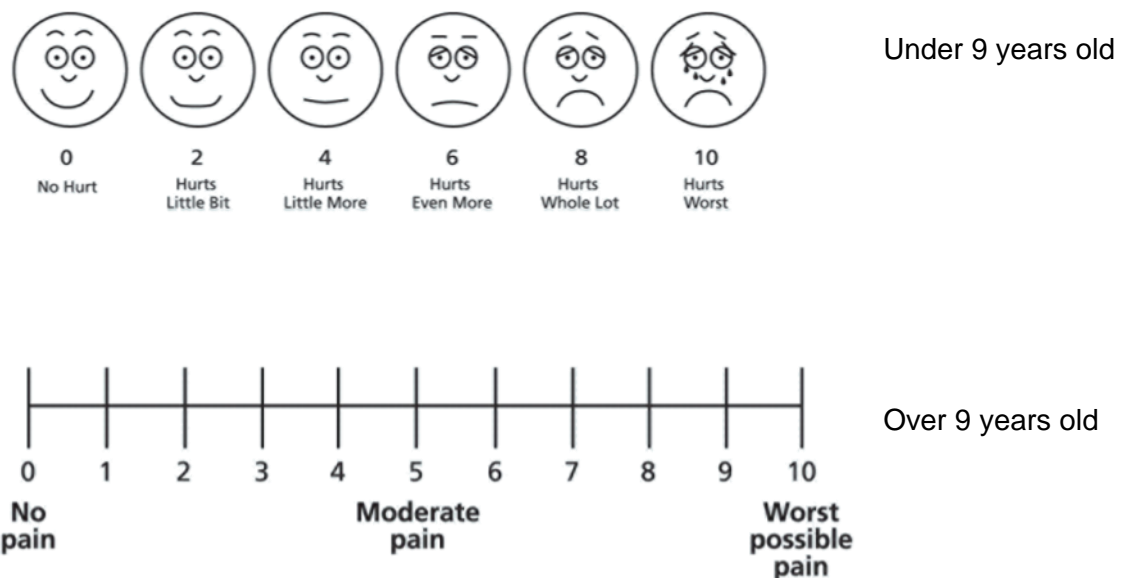
	WHO Grading	Joule setting /cm ²	Time sec/ cm ²
Grade 0	None	1	2
Grade 1	Soreness +/- erythema, no ulceration	2	4
Grade 2	Erythema, ulcers. Patients can swallow solid diet	4	8
Grade 3	Ulcers, extensive erythema. Patients cannot swallow solid diet	8	16
Grade 4	Oral mucositis to the extent that alimentation is not possible	16	32

Sites

Site	WHO scale	J/cm ²	Time sec/ cm ²	Pain Pre-	Pain Post
TOTAL Daily J/cm ²					

A pilot study completed in 2012 involving 15 patients demonstrated a statistically significant reduction in self-reported pain levels for children who had treatment with LLLT for mucositis. A larger, retrospective case note analysis was then undertaken, comprising 319 separate LLLT sessions in 39 patients undergoing LLLT treatment from January 2013 to July 2014. Paediatric patients with symptomatic oro-pharyngeal mucositis were included (aged 4-17 years old). Patients were excluded from this analysis if they were not able to communicate their pain score due to communication ability or age. The Diobeam 830 laser was used for all treatment, with administered dose (J/cm²) varying according to the severity of mucositis (WHO scale 1-4). The pain scores (0-10) were recorded prior to and immediately post-treatment on a visual/faces analogue scale (Figure 4), along with daily neutrophil counts. Neutrophils are the most common white blood cell and in this group of patients the counts vary greatly depending upon their stage of disease and type of treatment.

Figure 4



The results are summarised in the Tables 1- 4 below. The most common WHO grade for mucositis was Grade 3 (extensive erythema and ulceration and cannot swallow a solid diet). A statistically significant change in pain score was noted with the median change of pain score being 2 (P-value <0.0005). No statistically significant relationship between neutrophil count and pain response was found (P-value = 0.263), or indeed between pain response and type of tumour (blood/solid) (P-value = 0.121).

Results

WHO Grading	Number of patients
0	8
1	38
2	79
3	149
4	45

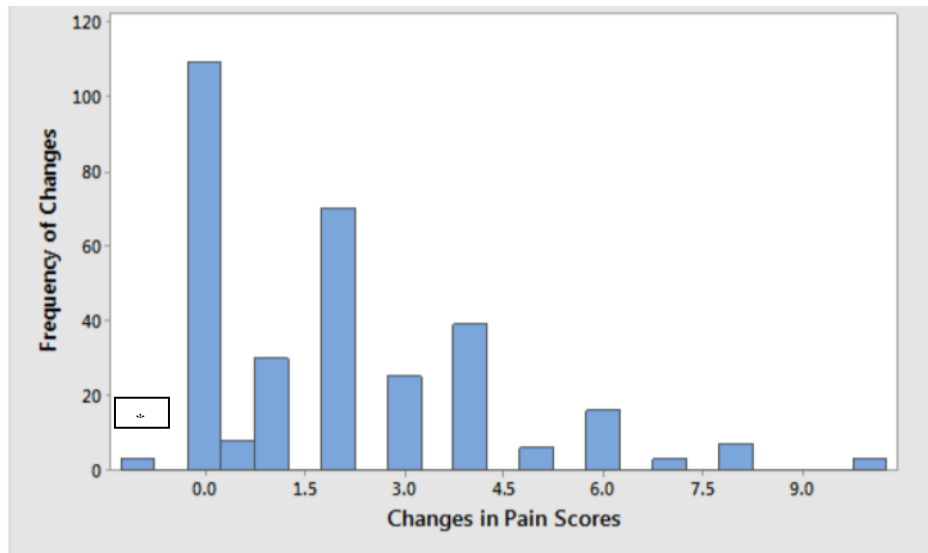
WHO Gradings pre-LLLT of the 319 episodes of treatment analysed

Type of Tumour	Number of Patients
Solid tumour	19
Blood borne	14
Inherited blood disorders	5
Solid and blood borne	1

Table 1: Summary of malignancies and WHO Gradings

Analysis of Pain Scores

Bar chart showing frequency of changes in pain scores



*One

patient reported an increase in pain score

Median change of pain score: 2 units

P-value <0.0005

For example: 8/10 pain score pre-treatment to 6/10 post-treatment

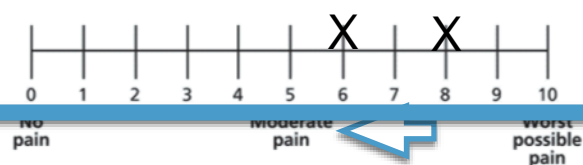
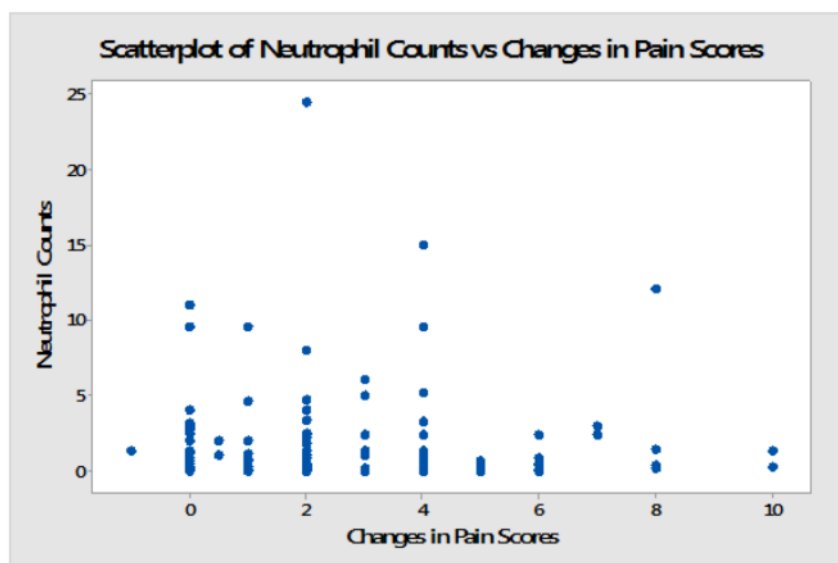
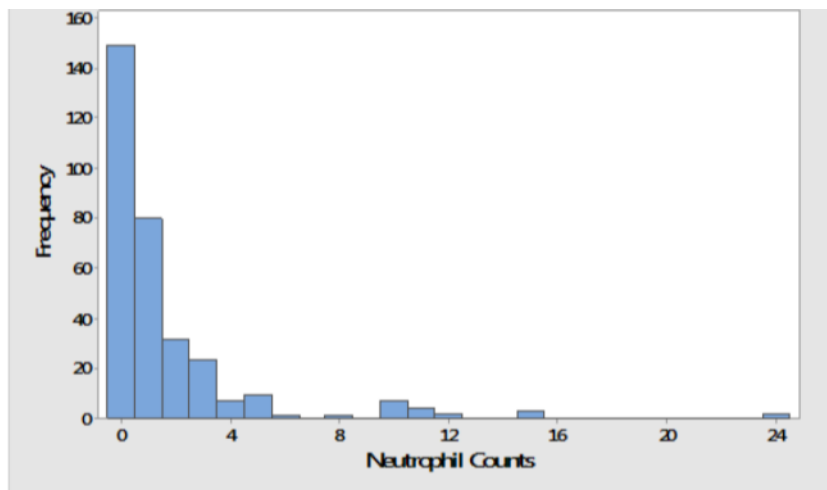


Table 2: Pain change results and number of patients with these changes

Neutrophil count of patients at the time of LLLT

Bar chart showing frequency of neutrophil counts



No statistical relationship between neutrophil count and pain response was found (P-value = 0.263)

Table 3: Summary of neutrophil counts at the time of LLLT treatment and comparison with changes in pain scores.

Pain Response versus Type of tumour

No statistical relationship between pain response and type of tumour, whether blood borne or solid tumour. (P-value = 0.121)

Table 4: Pain change and tumour results

Parents, staff and patients have embraced LLLT, finding it easy and beneficial, with regular requests for treatment and many positive reports. The ward and dental team work closely to ensure LLLT is available to those patients who need it and it has been widely accepted by the medical team as an effective management tool for mucositis.

Currently in Glasgow, due to resources and staff availability, LLLT is mainly provided when patients develop symptomatic mucositis. Our research has shown the ability to reduce pain in this patient group and this theoretically should reduce the need for narcotic analgesia and allow patients to maintain oral diets. This has been supported with one of the pain nurses in Glasgow stating that, 'The laser provides better pain relief than ketamine.' A patient explained when asked about LLLT 'my mouth was a 10/10, now it's a 0/10.' Figueiredo et al in 2013 ¹⁸ conducted a meta-analysis of the prophylactic use of LLLT to prevent mucositis and showed a significant positive effect. This research may extend our provision of LLLT, utilizing the lowest setting on the laser to provide this.

Conclusion

As with all new technologies, availability and cost naturally become less of a consideration over time. Access to LLLT across medical and dental services will no doubt improve, alongside cost effectiveness. A study by Bezinelli et al (2014) has already demonstrated that oncology patients with mucositis not treated with LLLT had 30% higher hospitalisation costs,¹⁹ an important finding for NHS hospitals under mounting financial pressures. Provision of LLLT is straightforward, quick and effective for many conditions. LLLT use within the specialist dental services continues to expand throughout the UK transforming the care we are able to routinely provide. Its use is also expanding in other areas of dental health and it will be interesting to see what the future holds for this remarkable technology.

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Figures

- Figure 1: Severe grade 4 oral mucositis
- Figure 2: Diobeam 830 Laser used in Glasgow
- Figure 3: The LLLT treatment sheet used in Glasgow
- Figure 4: Pain scales used in Glasgow

Tables 2 and 3

Graphs taken from statistical report by Johnathan Love B.Sc. 'Investigating the low level light therapy in paediatric patients suffering from chemotherapy induced oropharyngeal mucositis.'